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From perception threshold to ion channels - a computational study

Running title: Perception threshold to ion channels

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Abstract

Small surface area electrodes have successfully been used to preferentially activate cutaneous nociceptors, unlike conventional large area electrodes which preferentially activate large non-nociceptor fibers. Assessments of the strength duration relationship, threshold electrotonus, and slowly increasing pulse forms have displayed different perception thresholds between large and small surface electrodes, which may indicate different excitability properties of the activated cutaneous nerves. In the current study, the origin of the differences in perception thresholds between the two electrodes was investigated. It was hypothesized that different perception thresholds could be explained by the varying distributions of voltage-gated ion channels and by morphological differences between peripheral nerve endings of small and large fibers. A two-part computational model was developed in order to study activation of peripheral nerve fibers by different cutaneous electrodes. The first part of the model was a finite element model, which calculated the extracellular field delivered by the cutaneous electrodes. The second part of the model was a detailed multi-compartment model of an A δ -axon as well as an A β -axon. The axon models included a wide range of voltage-gated ion channels: Na_{TTXs}, Na_{TTXr}, Na_p, K_{dr}, K_M, K_A and HCN. The computational model reproduced the experimentally assessed perception thresholds for the three protocols, the strength duration relationship, the threshold electrotonus, and the slowly increasing pulse forms. The results support the hypothesis that voltage-gated ion channel distributions and morphology differences between small and large fibers were sufficient to explain the difference in perception thresholds between the two electrodes. In conclusion, assessments of perception thresholds using the three protocols may be an indirect measurement of the membrane excitability, and computational models may have the possibility to link voltage-gated ion channel activation to perception threshold measurements.

Introduction

The signaling pathway for our everyday perception of pain begins in the periphery through a select group of sensory neurons termed primary afferent nociceptors. Nociceptors are nerve fibers, which respond to damaging or potentially damaging stimuli and thus fulfill a protective purpose. During neuropathic conditions this protective function may be malfunctioning whereby the excitability of nociceptors has been shown to be increased (1). Neuropathic pain has a prevalence rate of 9.8 % which is approximately a third of the adult chronic pain patient group (2, 3). Both diagnosis and treatment of neuropathic pain is today inadequate and improved diagnostic tools as well as deeper understanding of the pathological mechanisms are needed. Increased excitability of the cell membrane has been identified in several neuropathic pain disorders (1) and several voltage gated ion channel abnormalities have been observed in patients with neuropathic pain disorders (4). Voltage-gated ion channels are suitable therapeutic targets for analgesic drugs since subtypes of voltage-gated ion channels are selectively expressed in nociceptors (5- 8) and due to their ability to govern the excitability of the nociceptors. Cutaneous electrical stimulation is particularly useful for studying activation of voltage-gated ion channels because the electrical current artificially activates the fibers through bypassing the sensory transduction at the nerve terminal. However, it is difficult to study the excitability of nociceptive fibers to electrical stimulation in isolation without coactivation of larger non-nociceptive fibers, since the electrical activation thresholds are lower for larger fibers compared to small fibers. To overcome this obstacle, small surface electrodes (Pin electrodes) have been developed (9-13) which generate a high current density in the epidermis where the small fibers terminate, unlike the large fibers which terminate in the dermis (14-17). As a result, the Pin electrodes have the potential to preferentially activate nociceptors at low stimulation intensities. Both objective estimations of the latency of evoked potentials (12-13) and human blink reflexes (10) as well as qualitative measurements of pain perception (9, 18), support that Pin electrodes preferentially activate

nociceptors. Non-invasive Pin electrodes preferentially activate the thinly myelinated A δ -fibers and intra-epidermal Pin electrodes have been reported to preferentially activate unmyelinated C-fibers as well as A δ -fibers (19).

The excitability of large fibers can be indirectly assessed by measuring the compound action potential unlike small fibers where the compound action potential has a slow velocity and therefore is spread out over time making it technically difficult to record. Therefore, our group has instead used perception thresholds as an indirect measurement of the excitability. In two recent studies, the perception thresholds for a concentric Pin electrode (cathode $\varnothing = 0.2$ mm) versus the conventional large surface patch electrode (Patch electrode, cathode 2 cm^2) were compared (18, 20). Both studies showed that perception thresholds differed between the two electrodes, which is a trivial consequence of the difference in electrode/tissue interface area. However, not only did the absolute values differ, but more interestingly, the differences were dependent on the duration and shape of the stimulation current which might indicate the influence of physiological mechanisms such as voltage-gated ion channels. In the Hennings et. al. study (20), two protocols to measure the perception thresholds were used, the strength duration protocol and the threshold electrotonus protocol. The strength duration protocol estimates the perception thresholds for different durations of rectangular pulses of the stimulation current. In that study, the strength duration time constant for the Patch electrode was significantly lower than the time constant for the Pin electrode. The second protocol, the threshold electrotonus, consists of a rectangular subthreshold prepulse followed by a short 1 ms rectangular pulse. The current of the short pulse is increased until the perception threshold is reached. The threshold electrotonus showed similar threshold reductions to subthreshold prepulses for depolarizing prepulses. However, for long hyperpolarizing prepulses, the Pin electrode demonstrated a significant increase in perception threshold compared to the Patch electrode. The study by Hugosdottir et. al. (18), showed that different shapes of slowly increasing stimulation currents alter the perception thresholds, particularly for long durations (50 ms) but only for the patch electrode. The novelty of these studies, is that they support that the perception threshold of cutaneous electrical stimulations can be used to indirectly assess excitability of axonal membranes.

In this study, a computational model has been developed in order to explain the differences in perception thresholds between the two fiber classes of sensory afferents. The hypothesis is that the main differences in perception threshold between the electrodes originate from the difference in morphology and voltage-gated ion channel distribution of peripheral nerves. For instance, TTX resistant sodium currents, both $\text{Na}_v1.8$ (Na_{TTXr}) and the persistent $\text{Na}_v1.9$ (Na_p), are preferentially expressed in nociceptors (6, 7, 21) and the current density of hyperpolarization-activated cyclic nucleotide-gated channels (HCN) is significantly positively correlated with the conduction velocity of these fibers (22). The expression and dynamics of potassium channels have also been shown to differ between fiber types (5) particularly the A-type potassium channel (K_A) (5, 8, 23). Therefore, the aim of this study was to investigate whether the difference in perception thresholds between the two electrodes could originate from the voltage-gated ion channel distribution as well as the morphological differences between peripheral nerve endings of large and small fibers.

Methods

A two-part computational model was developed in order to study activation of peripheral nerve fibers by cutaneous electrodes (see figure 1). The first part of the model is a finite elements model (COMSOL Multiphysics 6.3, Stockholm, Sweden) which calculates the electrical field delivered by a cutaneous electrode. The second part of the model is a detailed multi-compartment model (NEURON, Yale, USA (24)) of an A δ -axon as well as an A β -axon.

Finite element model for the electrical field

To calculate the electrical field along a nerve fiber the Mørch et. al. model (25) was further developed. The model consists of four rectangular skin layers: stratum corneum, epidermis, dermis and hypodermis. In each layer, the tissue was modeled as a linear and homogeneous medium. The electrical properties and thicknesses of the different layers were adopted from literature and are listed in table 1. Both a Pin electrode and a Patch electrode were implemented in the model. The full size of the model was 15x15 cm for the Pin model and 20x15 cm for the Patch model. The Pin electrode consisted of 15 steel pin cathodes placed in a circle and the anode was concentrically placed around the 15 pins (see figure 1A). The Pin electrode was designed with the same spatial dimensions as stated in the Lelic et. al. study (13). The spatial dimensions for the large surface area electrode cathode was 1.5x2 cm and the anode 4.5x6 cm. The anode was placed 12 cm from the cathode. The time dependent solution of the finite element model was calculated for rectangular pulses of the stimulation current (see Supporting material, figure S2). Since there was little effect from the tissue permittivity on the solution the steady state solution was used assuming the system is unchanging with regard to time. The following three boundary conditions were added to the model: the total current through the cathode was set to 1 mA, the total current through the anode was set to -1 mA and the hypodermis layer was grounded

underneath. The steady state solution of the electrical field was calculated by the Poisson equation:

$$-\nabla \cdot (\sigma \nabla V_{EF}) = 0 \quad \sigma \text{-Conductivity of the tissue, } V_{EF} \text{ electrical potential}$$

The iterative non-linear solver, Automatic Newton, was used with the maximum element size set to 4.2 mm and minimum 0.021 mm. To ensure convergence, both the mesh density and the size of the model were varied (see Supporting material, figure S1).

Tabel 1. The electrical properties and thicknesses of the different layers of the skin.

	Electrical conductivity (S/m)	Relative electrical permittivity	Thickness (μm)
Stratum Corneum	2x10 ⁻⁴ (26)	2x10 ³ (26)	20 (27, 28)
Epidermis	H:0.95 (29)	6.1x10 ⁴ (29)	82 (27, 28)
	V:0.15 (29)	2.6x10 ⁴ (29)	
Dermis	H:2.57 (29)	2.4x10 ⁵ (29)	1300 (30)
	V:1.62 (29)	9.6x10 ⁴ (29)	
Hypodermis	2x10 ⁻² (31, 32)	1x10 ³ (31, 32)	5000

Nerve fiber models

Morphology: To study how the extracellular field is affecting the activation of nerve fibers, two multi-compartmental nerve fiber models were developed, one of an Aδ-fiber axon (Ø: 3.5 μm) and another of an Aβ-fiber axon (Ø: 9 μm). In a previous study (33), a novel C-fiber model was developed, which was further developed into these two myelinated axon models for the purpose of this study. Since electrical stimulation artificially generates an action potential without activating the sensory receptors terminals, the process of sensory transduction within the sensory terminal was not included in the model. Since the sensory terminal was not included, the Aβ model is terminated by a node of Ranvier. Both ends of the model's termination conditions were

sealed end. To model a damped propagation of the action potential at the model termination, an additional passive segment was added at the end of the model.

The A δ axons lose their myelin when they cross the dermal-epidermal junction (16, 17) and become morphologically indistinguishable from unmyelinated fibers (15). The A δ model therefore consists of two parts: one myelinated axon located in the deeper tissues and one unmyelinated axon protruding into the epidermis. The morphology of the A δ -fiber was modeled as a straight line with a 45 degree angle from the middle of the epidermis to the middle of the hypodermis where it continues horizontally through the hypodermis. Two side branches were added to the main branch to generate a nerve fiber density of 0.58 fiber endings per millimeter (see figure 1C). The angle between the main branch and side branches was 90 degrees.

The A β model has a similar spatial location except that the nerve fiber terminates in the middle of the dermis (see figure 1C). Both nerve fiber models were placed centered underneath the cathode for the Patch electrode or underneath one of the pins of the Pin electrode. Six additional spatial positions of the nerve fiber model were evaluated. Both models were moved (40 μ m) in all three dimensions generating a total of seven models. The current needed to generate an action potential which propagates to the end model was calculated for each of the seven models.

As the nerve fiber enters the superficial layers of the skin, the length of the intermodal distance decreases, the myelination reduces and the length of the nodes of Ranvier is increased (15). Therefore, these parameters have been reduced by a factor of two from the morphology measured from fibers in deeper tissues (34, 35). All morphological parameters are listed in table 2. The electrical field calculated by the finite element model was added as the extracellular field to the axon models. The number of the compartments were 26,088 for the A δ model and 27,120 for the A β model, and the equations were solved using the variable time step method in NEURON.

Table 2. Parameters of the morphology for the axon models.

	A δ -fiber	A β -fiber	Reference
Diameter	3.5 μ m	9 μ m	

Nodal length	2 μm	2 μm	(15, 34)
Internodal length	400 μm	900 μm	(15, 36)
Juxtaparanodal length	5 μm	5 μm	(37)
Capacitance nodal	1 $\mu\text{F}/\text{cm}^2$	1 $\mu\text{F}/\text{cm}^2$	(38)
Capacitance myelin	0.012 $\mu\text{F}/\text{cm}^2$	0.0071 $\mu\text{F}/\text{cm}^2$	(38) $C=1/(\text{myelin sheet} + 1)$
Resting membrane potential	-60 mV	-60 mV	(39)
Number of myelin sheets	40	70	(15, 35)
Intra cellular resistance	130 Ωcm	130 Ωcm	
Branch Length	Main: 500 μm Side branch 1: 49 μm Side branch 2: 99 μm		(15)
Total model length	5.472 cm	4.994 cm	

Voltage-gated ion channel models: One of the main difficulties in the development of these axon models was to implement the densities of the ion channels. Larger fibers have been thoroughly researched but for smaller fibers, such as nociceptors, it is technically more difficult to perform intracellular recordings by patch clamp techniques. Therefore, most of the ion channel data is obtained from the dorsal root ganglion soma (DRG) and mainly from the smallest unmyelinated C-fibers. Details regarding the shape of the action potential and the relative influence of the underlying ion channel currents in A δ -fibers are not known. However, the major sodium current during an action potential in the DRG soma is Na_{TTXr} (40) most probably driven by the activation of Nav1.8 channels (6). Therefore, the Na_{TTXr} current is substantially larger than the Na_{TTXs} current in the computational model.

The ion channel currents were described by Hodgkin-Huxley-type ion channels. The myelinated fiber section of the A δ model has the same subtypes of ion channels as the A β -fiber model except for the subtypes of the Na_p channel and the K_A channel which is selectively expressed in nociceptors (8, 21). The HCN maximum conductance was additionally reduced by a factor of

two, since experimental patch clamp experiments have estimated that A δ neuron somas express half the current compared to the larger somas of A β neurons (22).

The literature supports large variations of the gating parameters (5), which may either be due to modulatory substances which are missing during patch clamp experiments, or to different compositions of the subtypes of the ion channels. Therefore, the gating parameters voltage dependencies were allowed to be shifted in order to fulfill the constraints. For the A β model, the ion channel dynamics for the delayed rectifier and persistent sodium channels were adopted from Jankelowitz et. al. model (41). With the original parameters for the delayed rectifier, the channel was activated prior to the action potential. The delayed rectifier ion channels' voltage dependence needed to be modified by a depolarizing shift of 40 mV in order for the channel not be activated before action potential generation, resulting in shifting the $V_{1/2}$ to -32 mV which is consistent with experimental observations (5). In this study, the fast TTX sensitive currents were denoted as Na_{TTXs} and the fast TTX resistant sodium current as Na_{TTXr} . The persistent sodium currents are denoted Na_p regardless of their sensitivity to TTX. The Jankelowitz et. al. model (41) did not include the slow inactivation gates for Na_{TTXs} channel (mainly $Na_v1.6$). Therefore, the ion channel dynamics were implemented from the Watanabe et. al. model (42) and the steady-state parameters were shifted 10 mV in order to generate an action potential which propagated to the proximal end of the axon model and generated depolarization larger than 10 mV. The ion channel dynamics for the M current and HCN were adopted from Tigerholm et. al. model (33).

For the A δ -fiber's unmyelinated nerve ending, the following three sodium channels were implemented: two TTX resistant currents (Na_{TTXr} and Na_p) and the Na_{TTXs} (mainly $Na_v1.7$). The steady state parameters and their time constants for the three sodium channels were taken from the Tigerholm et. al. model (33), as were the HCN current and the K_M current dynamics. The model of the K_A was adopted from Watanabe et. al model (42). The channel voltage dependency was modified by a 15 mV hyperpolarizing shift in order to get the model consistent with the threshold electrotonus measurements of the Pin electrode (20). For the A \square model, the K_A current steady-state parameters were set to: $V_m = -2.4$; $k_m = 14.2$; $V_h = 30.7$ and $k_h = 6.1$ (5). The time constant was implemented from the Tigerholm et al. study (33). The parameters for the voltage-gated ion channels are listed in table 3.

Table 3. Parameters for the axon models.

Aδ- model	Model Reference	Spatial location	Maximal Conductance (S/cm²)
Na_{TTXr} (Na_v1.8)	(33)	Unmyelinated nerve ending	$3.03 \cdot 10^{-2}$
Na_p (Na_v1.9)	(33)	Unmyelinated nerve ending and nodes of Ranvier	$2.96 \cdot 10^{-4}$ (Unmyelinated) $1.10 \cdot 10^{-3}$ (node)
Na_{TTXs} (Na_v1.6)	(42)	Nodes of Ranvier	$1.45 \cdot 10^{-1}$
Na_{TTXs} (mainly Na_v1.7)	(43)	Unmyelinated nerve ending	$1.27 \cdot 10^{-2}$
K_{dr}	(41)	Unmyelinated nerve ending and Juxtaparanode	$1.20 \cdot 10^{-3}$ (Unmyelinated) $4.80 \cdot 10^{-3}$ (Juxtaparanode)
K_A	(44)	Unmyelinated nerve ending and Juxtaparanode	$2.70 \cdot 10^{-3}$ (Unmyelinated) $3.00 \cdot 10^{-3}$ (Juxtaparanode)
K_M	(33)	Unmyelinated nerve ending and nodes of Ranvier	$2.62 \cdot 10^{-5}$ (Unmyelinated) $2.10 \cdot 10^{-3}$ (node)
HCN	(33)	Unmyelinated nerve ending and Juxtaparanode	$3.81 \cdot 10^{-5}$ (Unmyelinated) $1.52 \cdot 10^{-4}$ (Juxtaparanode)
Leak channel	(33)	Internode	$1.00 \cdot 10^{-7}$
Aβ- model			
Na_{TTXs} (Na_v1.6)	(42)	Nodes of Ranvier	$1.45 \cdot 10^{-1}$
Na_p	(41)	Nodes of Ranvier	$2.30 \cdot 10^{-5}$
K_A	(5)	Juxtaparanode	$3.90 \cdot 10^{-2}$
K_{Dr}	(41)	Juxtaparanode	$4.80 \cdot 10^{-3}$
HCN	(33)	Juxtaparanode	$3.04 \cdot 10^{-4}$
K_M	(33)	Nodes of Ranvier	$2.10 \cdot 10^{-3}$
Leak channel	(33)	Internode	$1.00 \cdot 10^{-7}$

Criteria used to set model parameters

The activation threshold was defined as the stimulation current required to generate an action potential which propagates to the end of the axon model. The maximum conductances for the voltage-gated ion channels were the unknown parameters in the two nerve fiber models. To constrain the model, three criteria were defined:

- 1) The A δ model should have a lower activation threshold than the A β model for the Pin electrode.
- 2) The A β model should have a lower activation threshold than the A δ model for the Patch electrode.
- 3) Three different protocols measuring the excitability of the membrane have been studied experimentally by recording the perception threshold using the Pin and the Patch electrode (18, 20). The model should be able to reproduce the differences in perception thresholds between the two electrodes for all three protocols (see figure 2). The three protocols are:

A. Strength duration curve, which is measuring the activation thresholds for different durations of rectangular pulses. The strength duration time-constant should be similar to the experimentally obtained values, which for the Pin electrode is $1060 \mu\text{s} \pm 690 \mu\text{s}$ and for the Patch electrode $580 \mu\text{s} \pm 160 \mu\text{s}$ (20). The time constants were estimated as the duration needed to generate an activation threshold of two-folded the rheobase (duration 16 ms). To increase the convergence of the axon model, the rise time for the electrical current was set to 0.025 ms.

B. Threshold electrotonus. First, the activation threshold for a rectangular 1 ms pulse was estimated. Thereafter, a prepulse was applied at a stimulation current corresponding to 20% of the initial activation threshold for the 1 ms pulse (i.e. before the 1 ms pulse was applied). The effect of the prepulse on the activation threshold for a 1 ms pulse was measured. The activation threshold was estimated for both hyperpolarizing and depolarizing prepulses.

C. Slowly increasing stimulation current. In the Hugossdottir et. al. study (18), the perception thresholds were compared for different shapes of slowly increasing stimulation

current for the two durations of 5 ms and 50 ms respectively. The different shapes and equations are listed in table 4.

Table 4. *The equation for the stimulation current for the slowly increasing stimulation protocol, see reference (18)*

Pulse form	Equation for the stimulation current	Label
Exponential increase	$I(t) = \begin{cases} \frac{I_s}{e^{\frac{T_s}{\tau}} - 1} \left(e^{\frac{t}{\tau}} - 1 \right), & 0 \leq t < T_s \\ I_s \cdot e^{\frac{-t}{\tau_{tr}}}, & T_s \leq t \leq T_{tr} \end{cases}$	<i>Exp</i>
Linear increase	$I(t) = \begin{cases} \frac{I_s}{T_s} \cdot t, & 0 \leq t < T_s \\ I_s \cdot e^{\frac{-t}{\tau_{tr}}}, & T_s \leq t \leq T_{tr} \end{cases}$	<i>Lin</i>
Bounded exponential	$I(t) = \begin{cases} \frac{I_s}{1 - e^{\frac{-T_s}{\tau}}} \left(1 - e^{\frac{-t}{\tau}} \right), & 0 \leq t < T_s \\ I_s \cdot e^{\frac{-t}{\tau_{tr}}}, & T_s \leq t \leq T_{tr} \end{cases}$	<i>ExpB</i>
Rectangular	$I(t) = I_s(t), \quad 0 \leq t < T_s$	<i>Rec</i>

I_s = stimulation current, T_s = stimulation duration $\tau = (T_s/2)$ = time constant.

Trailing phase: $T_{tr} = T_s \cdot 1.4$ and $\tau_{tr} = \tau/6.6$

Results

Activation of nerve fibers by cutaneous electrical stimulation

The distinction in spatial location of small and large fiber nerve endings is thought to generate the preferential activation of small fibers by the Pin electrode since the Pin electrodes have the potential to generate a high electrical current density in the most superficial skin layer (epidermis). To investigate this, the electrical field generated by the two electrodes are depicted in figure 3. The Pin electrodes generate a local and high electrical current which is highly restricted to the stratum corneum and epidermis, while the Patch electrode generates a larger spread of the electrical field resulting in a large depolarizing electrical field also in deep tissues.

To investigate how the electrical field from the electrodes is affecting the nerve fibers we compared the electrical field along two axons, one $A\beta$ and one $A\delta$, when a constant electrical stimulus of 1 mA was applied through the electrodes (figure 4). The Pin electrode generated a substantially higher electrical field in the epidermis (0.18 V) compared to the patch electrode (0.08 V). If the electrodes were moved on the surface of the skin, away from the tip of the axon model, the high electrical field in epidermis was largely reduced for the Pin electrode, indicating a small region of selective activation of small fibers for the Pin electrode (see Supporting material, Figure, S1). The Patch electrode was not as sensitive as the Pin electrode to movements of the electrode (see Supporting material, figure S1).

To study the action potential generation, a rectangular pulse with a duration of 2 ms was applied through the electrodes (see figure 5), leading to an action potential generated at the tip of the nerves. Although, when the $A\delta$ model was activated by the patch electrode, the action potential was generated in deeper tissue layers at the first nodes of Ranvier. In figure 5, changes in ion channel currents and membrane potential during an action potential are illustrated.

The slow ion channels' densities differ between the two models, particularly the K_M and the persistent sodium channels. For the $A\delta$ model, the persistent sodium is the dominating slow current with a factor of 5 times higher peak current than the K_M current during an action potential. For the $A\beta$ the dominating current is the K_M current which is a factor of 10 times larger during an action potential (see figure 5).

The nerve fiber model's activation thresholds for the three protocols

The stimulation current required to generate an action potential for the three different protocols was estimated for all seven spatial positions of the nerve fiber models. If the $A\delta$ model was moved down 40 μm closer to the dermis-epidermis junction, the current needed to activate the fiber model exceeded the value of the $A\beta$ model for the Pin electrode. Therefore, this model was not included when the average activation threshold was calculated for the $A\delta$ model. The strength duration curve protocol measures the activation threshold for different durations of rectangular pulses (see figure 6A). The $A\delta$ model showed a higher increase in activation threshold for shorter

pulses compared to the A β model. This is consistent with the experimental results showing a significantly higher time constant for the Pin electrode (computational model: $633 \pm 36 \mu\text{s}$, experimental result: $1060 \mu\text{s} \pm 690 \mu\text{s}$ (20)) than for the Patch electrode (computational model: $428 \pm 4 \mu\text{s}$, experimental result: $580 \mu\text{s} \pm 160 \mu\text{s}$ (20)). The strength duration curve is strongly influenced by the fast sodium current which generates the action potential. Subthreshold prepulses are effective to increase the influence of slower ion currents on the activation threshold. In the current study, the threshold electrotonus protocol was studied which examines the excitability changes related to long subthreshold rectangular prepulses (10-80 ms) – both depolarizing and hyperpolarizing prepulses. The A δ model and the A β model show similar threshold alterations to subthreshold depolarizing prepulses (see figure 6B). However, for long hyperpolarizing prepulses, the A δ model had a large increase in the activation threshold compared to the A β model. For 80 ms hyperpolarizing prepulses, the activation threshold was increased by 68 % in the A δ model but only by 25 % in A β model. This result is consistent with experimentally obtained measurements of the perception threshold where the only significant difference between the two electrodes' threshold electrotonus was for long hyperpolarizing prepulses where the Pin electrode had a significant increase in the perception threshold (61%) compared to the Patch electrode (27%) (20).

The last protocol measures the excitability for three different shapes of slowly increasing stimulation pulses: Exponential increase, Linear increase and Bounded exponential increase, each with the same durations of either 5 ms or 50 ms. For the 5 ms duration, both the A δ model and the A β model have a reduced activation threshold when the derivative of the initial phase of the electrical stimulus is high. For the longer 50 ms durations, the threshold is instead increased for the A β model from 0.73 (Exp 50 ms) to 0.82 (ExpB 50 ms) (see figure 6C). This is consistent with the experimental study which measures the perception thresholds for the Patch electrode (18). In the experimental study, the patch electrode increased the activation threshold from 0.76 (Exp 50 ms) to 0.94 (ExpB 50 ms) (see figure 6C). For the A δ model, the activation threshold is not increased for the longer 50 ms duration which is consistent with the activation threshold measured by the Pin electrode (see figure 6C). The rectangular pulse was chosen as a control and generated a lower threshold for the 50 ms duration compared to the slowly increasing pulses, which is likewise consistent with experimental observations (18).

Longer duration electrical stimuli may increase the preferential activation of small fibers

To study how selective the Pin electrode is for the strength duration and the slowly increasing stimulation current protocols, the activation thresholds of the different models were compared. For the strength duration protocol, the activation thresholds for rectangular pulses with different durations, are represented in figure 7. The activation threshold is considerably lower for the $A\beta$ model compared to the $A\delta$ model when the Patch electrode is activating both axonal models. However, this is not the case for the Pin electrode where the activation threshold is slightly lower for the $A\delta$ model for pulse durations longer than 0.4 ms. For durations shorter than 0.4 ms, the Pin electrode has a higher activation threshold for the $A\delta$ model, which indicates that the selective activation of small fibers is reduced for short durations of rectangular pulses.

The same behavior can be seen for the slowly increasing stimulation currents, where the difference in activation thresholds between the two models is larger for long durations compared to short durations for the Pin electrode (see figure 8C). The excitability change due to slowly increasing electrical stimuli is different between the two nerve models. The $A\delta$ model becomes more excitable when the membrane potential is increasing slowly and an action potential can be generated when the cell membrane is depolarized. For the $A\beta$ model, slow depolarization leads to inactivation of sodium channels and to the inability to generate an action potential when the membrane is too depolarized. This can be seen in figure 8B, where the action potential for the $A\beta$ model is not generated at the maximum depolarization level, but instead before, compared to the $A\delta$ model where the action potential is generated at the point of maximum depolarization of the cell membrane.

Varying the current densities of the voltage-gated ion channels

The computational model of peripheral nerve endings was sufficient to reproduce the differences in perception thresholds between the two fiber classes. To further study the underlying

mechanisms for the differences in perception threshold between the electrodes, the influence of different subtypes of ion channels was studied. The ion channel densities which are selectively expressed in the different classes of fibers were altered: the Na_{TTXs} channel, the Na_p channel, the K_A channel and the HCN channel. In small fibers, the action potential is generated by the TTX-resistant current (40) which is not the case in the node of Ranvier where the dominating sodium current is the Na_{TTXs} current (45). The Na_{TTXf} and the Na_{TTXs} have different dynamics (40) and thereby affect the excitability differently. In figure 9, the Na_{TTXs} current density in the $\text{A}\delta$ model was increased until an action potential was generated by the Na_{TTXs} current. This mainly reduced the activation threshold for short stimulus durations resulting in a largely reduced time constant for the strength duration curve (from 628 μs to 409 μs). Additionally, the strong influence of long hyperpolarizing subthreshold prepulses was reduced in the $\text{A}\delta$ model with higher Na_{TTXs} current density (from 60% to 30%). Moreover, for the $\text{A}\delta$ model both the strength duration curve and the threshold electrotonus became more similar to the perception threshold recorded for large fibers. For slowly increasing current stimulations, there was a minor change in the $\text{A}\delta$ model activation threshold with an increase in the Na_{TTXs} sensitive current.

If instead the Na_p channel's current density was reduced, the activation threshold for electrotonus and slowly increasing current stimulation was affected but the activation threshold for the strength duration curve remained unchanged (see figure 9). For instance, the strong effect of hyperpolarizing long prepulses (80 ms) in the $\text{A}\delta$ model was reduced from 60% to 46% and became more similar to the $\text{A}\beta$ model threshold (25%). Additionally, the low activation threshold for slowly increasing electrical stimulation (ExpB) was increased from 0.65 to 0.71 which is closer to the $\text{A}\beta$ model activation threshold (0.81).

When the current density for the subtype of Na_p channels expressed in large fibers was increased in the $\text{A}\beta$ model, the activation threshold for long slowly increasing current stimulation was reduced (from 0.81 to 0.74) and became more similar to the $\text{A}\delta$ model's activation threshold (0.65). A reduction of the HCN current density did not influence the activation threshold of the $\text{A}\beta$ model considerably.

Discussion

In a previous study, the excitability in an unmyelinated axon model was examined (25) and in the current study the model was further developed into an A δ model and an A β model and included a finite element model of the skin and two cutaneous electrodes. The origin of the differences in perception thresholds between two electrodes have been studied, and the hypothesis was that the main difference could be explained by different distributions of voltage-gated ion channels and along with differences in morphology between peripheral nerve of small and large fibers. The voltage-gated ion channel densities and morphological differences between the two fiber classes were sufficient to reproduce the main difference in perception threshold between the two electrodes; particularly the voltage-gated ion channels play a pivotal role.

Computational models of axons and their clinical implications

Even though the field of computational models of central and peripheral nerves arose from Hodgkin's and Huxley's squid axon model (46) the number of existing detailed computational models of peripheral small fibers is surprisingly low. Since it is technically difficult to perform intracellular recordings on axons, particularly on thin fibers, there is a lack of data regarding ion channel densities and dynamics. One of the most detailed computational models of a nociceptor is an unmyelinated axon model (33, 47, 48) which was constrained by latency measurements recorded by microneurography. The computational model developed in the present study is a combination between the C fiber model and the A δ /A β model developed in the Mørch et. al. study (25). The current computational model combines a computational model of the skin to estimate the extracellular potential with a detailed nerve fiber model with a wide range of voltage-gated ion channels. Existing spaced-clamped models, which lack spatial distribution, have been able to link threshold tracking measurements of compound motor and sensory action potentials to subtypes of voltage-gated ion channels (49-53). The novelty of the present computational model is that it can link perception threshold measurements to voltage-gated ion channel activation in both an A δ model and an A β model. Furthermore, spaced-clamped models of large fibers have

been successfully used to identify abnormalities in voltage-gated ion channel densities during pathological conditions (41, 54). The current computational model may be used in a similar fashion. Since perception threshold tracking is a non-invasive and inexpensive method, the model could be a useful tool for understanding the mechanisms of painful peripheral neuropathy, particularly due to channelopathies. Increased excitability has been identified as an underlying mechanism in many peripheral neuropathies (1). However, the underlying ionic current or passive property alterations resulting in such increased excitability is essentially unknown. For instance, the identification of a specific voltage-gated ion channel or groups of ion channels contributing to the excitability, could be used as a powerful diagnostic tool as well as potentially provide important guidance to the development of novel analgesic drugs. Different subtypes of ion channels affect different aspects of membrane excitability. For instance, slowly activating ion channels mainly affect the excitability when long current simulation is applied, while rapidly activating channels have strong influence on excitability of short current stimulations (see figure 9). Assuming that a pathological regulation (up or down) of a voltage-gated ion channel has occurred, would it be possible to distinguish which subtypes of ion channels have been altered by an intelligent measure of the perception threshold? By means of example: The three ion channels, the persistent sodium channel, the Na_{TTXs} channel and the K_A potassium channel, have all been implicated with peripheral neuropathy (55-57). If the Na_{TTXs} current is increased, the activation threshold for small rectangular pulses will be significantly reduced, while an increase of the persistent sodium current would mainly affect excitability for slowly increasing pulses or subthreshold prepulses (see figure 8). A reduction of the K_A current would mainly affect the excitability for slowly increasing pulses. All three alterations of ion channels should therefore theoretically be possible to identify by perception threshold measurements. This indicates also that computational models may be used as a diagnostic tool to identify ion channelopathies related to small fiber neuropathy.

The mechanism underlying the differences in perception threshold between the electrodes

The excitability in large fibers can be studied by threshold tracking techniques measuring the compound muscle action potential (58). However, the small nociceptive fibers' compound action

potential is technically difficult to assess. Our research group has therefore previously used the perception threshold measurements instead of the compound action potential as an indirect measurement of the excitability in the small fibers (20, 18). The disadvantage of using the perception threshold instead of the compound action potential is the influence of the central nervous system on the measurements, such as habituation, expectation and attention. Fortunately, the main effect on the perception threshold seems to originate from peripheral nerve mechanisms since excitability protocols, such as the threshold electrotonus and strength duration curve, display very similar behavior regardless of whether the measurements relate to the perception threshold or the compound action potential (58, 20). Perception thresholds may therefore be used as an indirect measurement of the membrane excitability of nerve fibers (20). Moreover, the computational model could reproduce the experimentally measured differences in perception thresholds between the two electrodes. When the densities of the ion channels were altered, this had a strong effect on the activation threshold of both models. Interestingly, when the Na_{TTXs} , which is highly expressed in nodes of Ranvier (45), was increased in the $\text{A}\delta$ model, the strength duration relation chronaxie was reduced from 628 μs to 409 μs approaching the value for the $\text{A}\beta$ model 428 μs . Also the large increase of the activation threshold for hyperpolarizing prepulses reduced from 60% to 30%, which is within range of the experimentally measured value of the patch electrode. This result implicates that the major difference between the strength duration relationship and the threshold electrotonus of the two electrodes could be explained by the expression of Na_{TTXs} .

Preferential activation of small nerve fibers by cutaneous electrodes

Preferential activation of small fibers by cutaneous electrodes has been performed by several different types of Pin electrodes, all with the common feature of a small cathode (9-13). Both objective measurements, such as conducting velocity measurements (12, 13, 19) and reflex measurements (10), as well as subjective measurements, such as the quality of the perception (18, 9) support the preferential activation of small fibers by a small area cathode. In our previous computational study (25), the results support that the Pin electrode has the potential to generate a high electrical field in the epidermis where the free nerve endings of small fibers are located. In

this study, a more detailed computational model was used, showing that this explanation seems to be valid. Even though the Pin electrodes preferentially activate nociceptors, the non-invasive pin electrodes mainly activate A δ fibers and not C-fibers. The ability to increase the preferential activation of small fibers would be beneficial for studying C-fibers. The results from the computational model predict that the small fibers' nerve endings need to be located close to the pins of the electrode in order for the electrode to preferentially activate these fibers. Multiple pins on the electrode may therefore be preferable to increase the amount of small fibers being activated concurrently which might be needed for the assessment of the perception threshold. Additionally, the results from the model predict that a longer duration of the electrical current, particularly slowly increasing electrical currents, may increase the preferential activation of nociceptors. As long as up to 50 ms durations of slowly increasing electrical input has been proposed to be used in order to increase the preferential activation of nociceptors (60, 18). The results from the present computational models support these experimental results and suggest that the underlying mechanism generating the increase in preferential activation is a selective expression of specific ion channels in nociceptors. To generate the reduced perception threshold of nociceptors for long slowly increasing electrical stimulus, the model predicted that both the Na_{TTXs} current as well as the K_M current should be minimally expressed. This result is consistent with the excitability alteration due to long ramp pulses measured in large fibers, where the reduced excitability for long durations of slowly increasing electrical input is generated by inactivation of sodium channels and the K_M current (58). Small fibers may be less sensitive to depolarization of the membrane since the Nav1.8, selectively expressed in nociceptors, inactivates at more depolarized membrane potential than the Na_{TTXs} channels expressed at the node of Ranvier (40).

Limitations of this study

In the finite element model, several features of the skin were omitted in order to reduce the computational complexity. For instance, the junctions between the layers were modeled as horizontal planes where the skin layers represented the average features of the skin. The skin is not an homogenous tissue but are innervated by sweat glands and hair cells. For instance, when

the electrode is moved over the skin surface the absolute value of the perception threshold is altered due to for instance the proximity to a nerve ending or a sweat gland. The heterogeneous composition of the skin and the electrode-skin interface has a strong influence on the absolute value of the perception threshold but less so on the relative difference in perception threshold for electrical stimulation in the range between 0.2 ms and 50 ms. For such reasons, the normalized perception threshold were compared to reduce the effect of heterogeneous skin composition. The skin's dynamic effect on the electrode current is small and has therefore been omitted (Supporting material, figure S2).

Even though the computational model includes seven voltage-gated ion channels, the model does not include any calcium channels which play an important role in regulating the excitability changes in nerve fibers. Finally, the major limitation to this study is the fact that the effect from the central nervous system on the perception threshold was omitted.

Conclusions

Since a computational model of peripheral nerve endings could successfully reproduce the experimental results from recorded perception threshold measurements, the hypothesis that peripheral membrane excitability as well as morphology contributes significantly to the perception threshold was supported.

Additionally, the results from this study indicate that a computational model could have the potential to link perception threshold tracking to the activation of specific ion channel subtypes, for instance, Na_{TTXs} , Na_{TTXr} and Na_{p} . Furthermore, the two computational models could be used to investigate how to further increase preferential activation of nociceptors by comparing the activation threshold differences.

Author Contributions

JT, AHP, CDM and OKA conceived and designed the research. JT carried out all simulations and analyzed the data. JT, AHP, CDM and OKA wrote the article.

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References

1. Serra. J, Bostock H, Solà R, Aleu J, García E, Cokic B, Navarro X, C. Quiles, 2012, Microneurographic identification of spontaneous activity in C-nociceptors in neuropathic pain states in humans and rats, *Pain*, 153:42-55.
2. Yawn B.P., Wollan P.C., Weingarten T.N., Watson J.C., Hooten W.M., Melton L.J., 2009, The prevalence of neuropathic pain: Clinical evaluation compared with screening tools in a community population, *Pain Medicine.*,10:586–593.
3. Portenoy R.K, Ugarte C., Fuller I, Haas G, 2004, Population-based survey of pain in the United States: differences among white, African American, and Hispanic subjects., *Journal of Pain.*, 5:317-28.
4. Waxman S.G. and Zamponi G.W, 2014, Regulating excitability of peripheral afferents: emerging ion channel targets, *Nature Neuroscience*, 17:153–163.
5. Gold M.S, Shuster M.J., Levine D.J, 1996, Characterization of six voltage-gated K⁺ currents in adult rat sensory neurons., *Journal of Neurophysiology.*, 75:2629-2646.
6. Akopian A.N., Sivilotti L., Wood J.N., 1996, A tetrodotoxin-resistant voltage-gated sodium channel expressed by sensory neurons., *Nature*, 379:257–262.
7. Djouhri L., Fang X., Okuse K., Wood J. N., Berry C. M., Lawson S. N., 2003, The TTX-resistant sodium channel Nav1.8 (SNS/PN3): expression and correlation with membrane properties in rat nociceptive primary afferent neurons., *Journal of Physiology.*, 550:739-752.

8. Phuket T.R and Covarrubias M., 2009, Kv4 Channels Underlie the Subthreshold-Operating A-type K⁺-current in Nociceptive Dorsal Root Ganglion Neurons, *Frontiers in Molecular Neuroscience.*, vol. 2:3.
9. Bromm B. and Meier W., 1984, The intracutaneous stimulus – a new pain model for algometric studies, *Methods Find Exp Clin Pharmacol*, 6:405–410.
10. Kaube H., Katsarava Z., Käufer T., Diener H., Ellrich J., 2000, A new method to increase nociception specificity of the human blink reflex., *Clinical Neurophysiology*, 111: 413-416.
11. Klein T., Magerl W., Hopf H., Sandkühler J., Treede R.D, 2004, Perceptual Correlates of Nociceptive Long-Term Potentiation and Long-Term Depression in Humans, *Journal of Neuroscience*, 24:964 –971.
12. Inui K, Tran D. T., Hoshiyama M., Kakigi R., 2002, Preferential stimulation of A δ fibers by intra-epidermal needle electrode in humans, *Pain*, 96:247–252
13. Lelic D., Mørch C.M., Hennings K., Andersen O.K., Drewes A.M., 2012, Differences in perception and brain activation following stimulation by large versus small area cutaneous surface electrodes, *European journal of pain*, 16:827-837.
14. Hilliges M, Wang L., Johansson O., 1995, Ultrastructural evidence for nerve fibers within all vital layers of the human epidermis, *Journal of Investigative Dermatology.*, 104:134-137.
15. Provitera V., Nolano M., Pagano A., Caporaso G., Stancanelli A., Santoro L., 2007, Myelinated nerve endings in human skin, *Muscle Nerve*, 35:767-775.
16. Ebenezer G.J., McArthur J.C, Thomas D., Murinson B., Hauer P., Polydefkis M., Griffin J.W, 2007, Denervation of skin in neuropathies: the sequence of axonal and Schwann cell changes in skin biopsies, *Brain.* , 130:2703-2714.
17. Myers I. M., Peltier A. C., Li J., 2013, Evaluating dermal myelinated nerve fibers in skin biopsy, *Muscle Nerve*, 47:1–11.
18. Hugosdottir R., Mørch C. D., Andersen O. K., Helgason T., Arendt-Nielsen L., 2017, Evaluating the ability of non-rectangular electrical pulse forms to preferentially activate nociceptive fibers by comparing perception thresholds, *Scandinavian Journal of Pain*, vol. 16:175.

- 19 Otsuru N., Inui K., Yamashiro K., Miyazaki T., Ohsawa I., Takeshima Y., Kakigi R., 2009,
 . Selective Stimulation of C Fibers by an Intra-Epidermal Needle Electrode in Humans, The
 Open Pain Journal, 2:53-56.
- 20 Hennings .K, Frahm K.S., Petrini .L, Andersen O.K., Arendt-Niels L., Mørch C., 2017,
 . Membrane properties in small cutaneous nerve fibers in humans., Muscle Nerve, 55:195-
 201.
- 21 Dib-Hajj, S. D. , Tyrrell L., Cummins T. R., Black J.A., Wood P. M, Waxman S. G., 1999,
 . Two tetrodotoxin-resistant sodium channels in human dorsal root ganglion neurons., FEBS
 Let, 462:117-120,
- 22 Gao L. L., McMullan S., Djouhri L., Acosta C., Harper A. A., Lawson S. N., 2012,
 . Expression and properties of hyperpolarization-activated current in rat dorsal root ganglion
 neurons with known sensory function., Journal of Physiology, 590: 4691-4705
- 23 Sarria I., Ling J., Gu J.G., 2012, Thermal sensitivity of voltage-gated Na⁺ channels and A-
 . type K⁺ channels contributes to somatosensory neuron excitability at cooling temperatures, J
 Neurochem, 122: 1145–1154.
- 24 Hines M. L. and. Carnevale N. T, Hines Carnevale, 1997, NEURON simulation
 . environment, Neural Computation, 9:1179-1209.
- 25 Mørch D. C., Hennings K., Andersen O. K., 2011, Estimating nerve excitation thresholds to
 . cutaneous electrical stimulation by finite element modeling combined with a stochastic
 branching nerve fiber model, Medical & Biological Engineering & Computing, 49:385–395.
- 26 Yamamoto T., Yamamoto Y., 1976, Electrical properties of the epidermal stratum corneum,
 . Medical & Biological Engineering , 14:151-158.
- 27 Sandby-Møller J., Poulsen T., Wulf H.C., 2003, Epidermal thickness at different body sites:
 . relationship to age, gender, pigmentation, blood content, skin type and smoking habits, Acta
 Dermato Venereologica., 86:410-413,.
- 28 Neerken S., Lucassen G. W., Bisschop M. A., Lenderink E. and Nuijs T., 2004,
 . Characterization of age-related effects in human skin: A comparative study that applies
 confocal laser scanning microscopy and optical coherence tomography., Journal of
 Biomedical Optics, vol. 9:274-281

- 29 Taverniera A., Dierickxb M., Hinsenkampa M., 1993, Tensors of dielectric permittivity and conductivity of in vitro human dermis and epidermis, *Bioelectrochemistry and Bioenergetics*, 30:65-72,
- 30 Krackowizer P. and Brenner E., 2008, Thickness of the human skin, *Phlebologie*, 37: 83-92.
- 31 Gabriel S., Lau R. W., Gabriel C., 1996, The dielectric properties of biological tissues: II. Measurements in the frequency range 10 Hz to 20 GHz, *Physics in medicine and biology*, 41:2251–2269.
- 32 Gabriel C., Peyman A., Grant E. H, 2009, Electrical conductivity of tissue at frequencies below 1 MHz, *Physics in medicine and biology*, 54:4863–4878.
- 33 Tigerholm J.*, Petersson M. E*, Obreja O., Lampert A., Carr R. W., Schmelz M., Fransén E., 2014, Modeling activity-dependent changes of axonal spike conduction in primary afferent C-nociceptors., *Journal of Neurophysiology*, 111:1721-35, * contributed equally.
- 34 Berthold M. and Rydmark C.H., 1983 Electron microscopic serial section analysis of nodes of Ranvier in lumbar spinal roots of the cat: A morphometric study of nodal compartments in fibres of different sizes, *Journal of Neurocytology*, 12:537–565.
- 35 Berthold C. H., Nilsson I., Rydmark M., 1983, Axon diameter and myelin sheath thickness in nerve fibres of the ventral spinal root of the seventh lumbar nerve of the adult and developing cat., *Journal of Anatomy*, 136:483-508.
- 36 Nilsson I. and Berthold C. H., 1988, Axon classes and internodal growth in the ventral spinal root L7 of adult and developing cats., *Journal of Anatomy*, 156,:71-96.
- 37 Poliak S., Salomon D., Elhanany H., Sabanay H., Kiernan B., Pevny L., Stewart C. L., Xu X., Chiu S. Y., Shrager P., Furley A. J., Peles E., 2003, Juxtaparanodal clustering of Shaker-like K⁺ channels in myelinated axons depends on Caspr2 and TAG-1., *Journal of Cell Biology*, 162:1149-1160.
- 38 Amir R. and Devor M., 2003, Electrical excitability of the soma of sensory neurons is required for spike invasion of the soma, but not for through-conduction., *Biophysical Journal*, 84:2181-2191.

- 39 Fang X., McMullan S., Lawson S. N., Djouhri L., 2005, Electrophysiological differences between nociceptive and non-nociceptive dorsal root ganglion neurones in the rat in vivo, *Journal of Physiology*, vol. 565, p. 927–943.
- 40 Blair N.T. and Bean B. P., 2002, Roles of tetrodotoxin (TTX)-sensitive Na⁺ current, TTX-resistant Na⁺ current, and Ca²⁺ current in the action potentials of nociceptive sensory neurons., *Journal of Neuroscience*., 22:10277-10290.
- 41 Jankelowitz S. K., Howells J., Burke D., 2007, Plasticity of inwardly rectifying conductances following a corticospinal lesion in human subjects., *Journal of Physiology*, 581:927-940.
- 42 Watanabe S., Hoffman D. A., Migliore M., Johnston D., 2002, Dendritic K⁺ channels contribute to spike-timing dependent long-term potentiation in hippocampal pyramidal neurons., *Proc Natl Acad Sci U S A*, 99:8366-8371.
- 43 Sheets P. L., Jackson J. O., Waxman S. G., Dib-Hajj S. D., 2007, Cummins T. R., A Nav1.7 channel mutation associated with hereditary erythromelalgia contributes to neuronal hyperexcitability and displays reduced lidocaine sensitivity, *Journal Physiology*, 581:1019–1031,
- 44 Migliore M., Hoffman D. A., Magee J. C., Johnston D., 1999, Role of an A-type K⁺ conductance in the back-propagation of action potentials in the dendrites of hippocampal pyramidal neurons., *Journal Computational Neuroscience*, 7:5-15.
- 45 Caldwell J. H., Schaller K. L., Lasher R. S., Peles E. Levinson S. R., 2000, Sodium channel Nav1. 6 is localized at nodes of ranvier, dendrites, and synapses., *Proc Natl Acad Sci U S A*., 97:5616–5620.
- 46 Hodgkin A. L. and Huxley A. F., 1952, A quantitative description of membrane current and its application to conduction and excitation in nerve, *Journal Physiology*, 117:500–544.
- 47 Tigerholm J.*, Petersson M. E.*, Obreja O., Eberhardt E., Namer B., Weidner C., Lampert A., Carr R. W., Schmelz M., Fransén E., 2015, C-fiber recovery cycle supernormality depends on ion concentration and ion channel permeability, *Biophysical Journal*, 108:1057-1071, * contributed equally.

- 48 Petersson M. E., Obreja O., Lampert A., Carr R.W., Schmelz M., Fransén E., Differential axonal conduction patterns of mechano-sensitive and mechano-insensitive nociceptors--a combined experimental and modelling study. 2014, PLoS One, 9.
- 49 Bostock H., Baker M., Reid G., 1991, Changes in excitability of human motor axons underlying post-ischaemic fasciculations: evidence for two stable states., *Journal of Physiology.*, 441:537–557.
- 50 Kiernan M. C., Isbister G. K., Lin C. S., Burke D., Bostock H., 2005, Acute tetrodotoxin-induced neurotoxicity after ingestion of puffer fish., *Annals of Neurology*, 57:339–348.
- 51 Howells J., Czesnik D., Trevillion L., Burke D., 2013, Excitability and the safety margin in human axons during hyperthermia, *Journal Physiology*, 591:3063–3080.
- 52 Trevillion L., Howells J., Bostock H., Burke D., 2010, Properties of low-threshold motor axons in the human median nerve, *Journal Physiology*, 588:2503–2515.
- 53 Howells J., Bostock H., Burke D., 2016, Accommodation to hyperpolarization of human axons assessed in the frequency domain, *Journal Neurophysiology*, 116:322–335.
- 54 Horn S., Quasthoff S., Grafe P., Bostock H., Renner R, Schrank B., 1996, Abnormal axonal inward rectification in diabetic neuropathy, *Muscle and nerve*, 19:1268-1272.
- 55 Dib-Hajj S. D., Black J. A., Waxman S. G., 2015, NaV1.9: a sodium channel linked to human pain., *Nature Reviews Neuroscience*, 16:511-519.
- 56 Hoeijmakers J. G., Faber C. G., Merkies I. S., Waxman S.G, 2015, Painful peripheral neuropathy and sodium channel mutations., *Neuroscience Letters*, 596:51-59.
- 57 Cao X. H., Byun H. S., Chen S. R., Cai Y. Q., Pan H.L., 2010, Reduction in voltage-gated K⁺ channel activity in primary sensory neurons in painful diabetic neuropathy: role of brain-derived neurotrophic factor., *Journal of Neurochemistry*, 114:1460–1475.
- 58 Bostock H., Cikurel K., Burke D., 1998, Threshold tracking techniques in the study of human peripheral nerve, *Muscle & Nerve*, p. 137–158, 1998.
- 59 Inui K. and Kakigi K., 2012, Pain perception in humans: use of intraepidermal electrical stimulation., *Journal of Neurology, Neurosurgery, Psychiatry*, 83: 551-556.

60 Jonas R., Namer B., Stockinger L., Chisholm K., Schnakenberg M., Landmann G.,
Kucharczyk M., Konrad C., Schmidt R., Carr R., McMahon S., Schmelz M., Rukwied R.,
2018, Tuning in C-nociceptors to reveal mechanisms in chronic neuropathic pain., *Annals of
Neurology*, Epub ahead of print.

FIGURE 1. Computational model design. *A two part computational model has been developed. The first part of the model calculates the electrical field generated by the two electrodes: a Pin electrode (A) or a Patch electrode (B). The skin model consists of four rectangular skin layers (C). The second part of the model consists of two axon models ($A\delta$ and $A\beta$) with the spatial location in the skin model illustrated in (C). The morphology of the myelinated axon sections consists of three parts: node of Ranvier, juxtaparanode and internode (D). The figures are not drawn to scale.*

FIGURE 2. The normalized time dependent stimulation currents for the three protocols. *A. The 6 rectangular shaped pulses used for the strength duration curve. B. An example of the subthreshold prepulse used during the threshold electrotonus protocol. C. The four different shapes of slowly increasing stimulation current used for the slowly increasing stimulation current protocol.*

FIGURE 3. The electrical field generated by the two electrodes. *A constant 1 mA stimulation current was applied through the electrode and the electrical field was calculated by the finite element method. A. The electrical field generated underneath the cathode of the Patch electrode. B. The electrical field generated underneath one of the Pin cathodes. C. Enlargement of the electrical field generated underneath one of the pin cathodes. The spatial location of the two axon models are illustrated by the two black lines. Note the spatially localized and the large depolarization generated by the Pin electrode in epidermis primarily innervated by nociceptive nerve fibers.*

FIGURE 4. The electrical field generated by the electrodes along the axons. *The figure to the left depicts the Pin electrode simulation and the figure to the right corresponds to the Patch electrode stimulation. The figure illustrates the electrical field generated by the two*

electrodes along the two axon models when a 1 mA continuous pulse was applied through the electrodes. Note the large difference between the electrical fields at the tip of the two axons when the stimulation current was applied through the Pin electrode. The lower two figures illustrate the electrical field generated along the branches for the $A\delta$ model.

FIGURE 5. Ion channel currents during an action potential. The figures to the left correspond to the simulation with the Pin electrode ($A\delta$ model) and the figures to the right correspond to the simulation with the Patch electrode ($A\beta$ model). Following a single rectangular pulse shaped stimulation current (0.4 mA) with a duration of 2 ms, an action potential was generated. A. Membrane potential and ionic currents recorded at the tip of the axon for both the $A\delta$ and $A\beta$ model. B. The large ion channel currents. C. The small ion channel currents. In the $A\beta$ model, the current density for the potassium current is low for both the K_A current and K_{Dr} current since the combined area of the juxtaparanode is 5 times larger than the node of Ranvier.

FIGURE 6. The activation threshold for the three protocols. The figures to the left correspond to the average activation thresholds estimated by the computational models and the figures to the right correspond to experimental data (The data was given from the authors on request (20, 18)). The activation threshold was defined as the stimulation current needed to generate an action potential which propagated to the end of the axon model. The average activation thresholds for different spatial locations is illustrated in the figure to the right. The error bars represent the standard deviation for the computational results and standard error for the experimental results.

For abbreviation description for the pulse shapes (Exp, Lin, ExpB and Rec), see table 4.

FIGURE 7. The strength duration curve for the Pin and the Patch electrode.

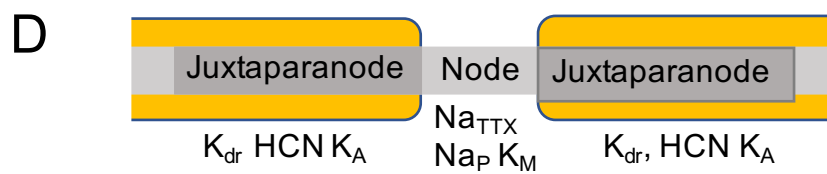
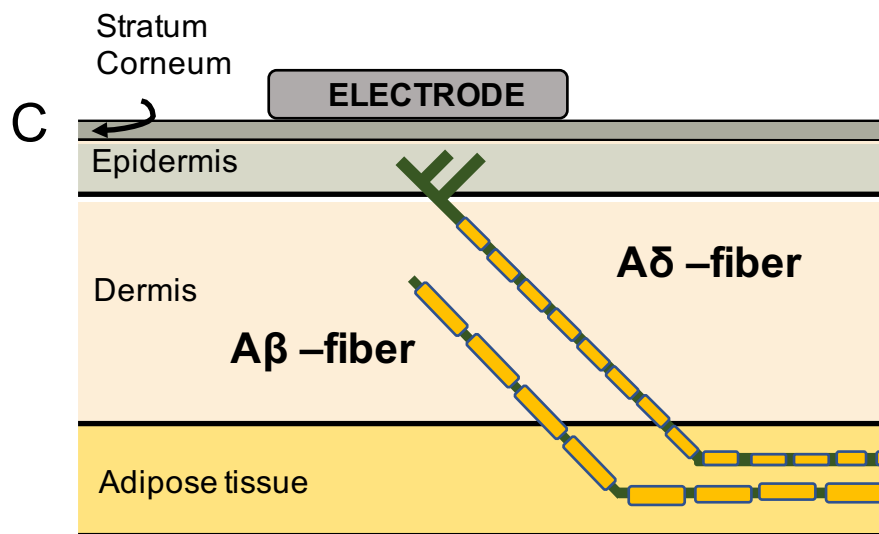
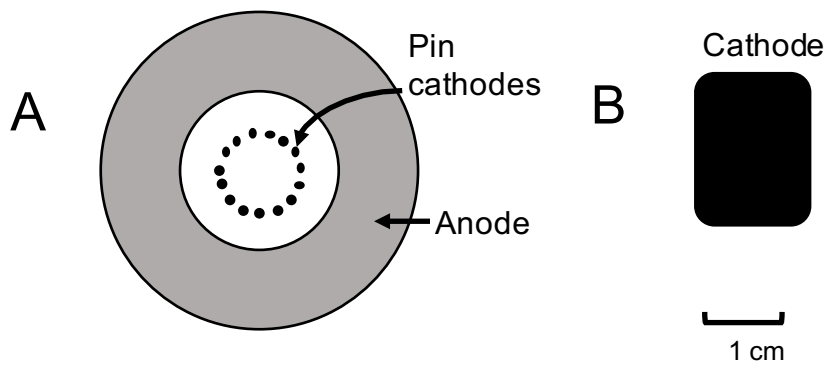
A. The stimulation current through the electrodes. B. The membrane potential corresponding to an extracellular field alteration when applying the current seen in A. through the electrode. The nerve fiber model was placed underneath the electrode in the middle of the epidermis ($A\delta$ model) or the dermis ($A\beta$ model). C. The average current needed to generate

an action potential for different spatial location is illustrated in the figure. The error bars represent the standard deviation.

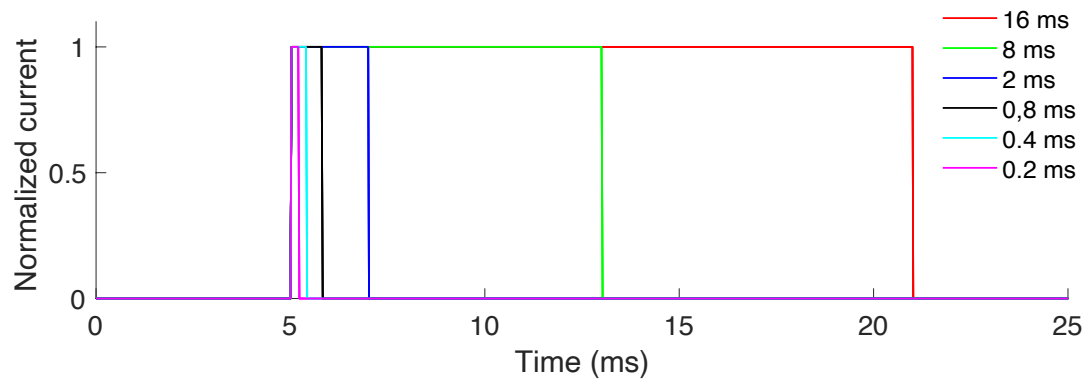
FIGURE 8. The effect of slowly increasing electrical stimulation on the preferential activation of small fibers. A. The stimulation current needed to generate an action potential for different shapes and durations of slowly increasing stimulation current. B. The membrane potential corresponding to an extracellular field alteration when applying the electrical current with the ExpB shape (see table 4). The nerve fiber model was placed underneath the electrode in the middle of the epidermis ($A\delta$ model) or the dermis ($A\beta$ model). C. The average current needed to generate an action potential for different spatial location is illustrated in the figure. The error bars represent the standard deviation.

FIGURE 9. Varying the Ion channel current densities

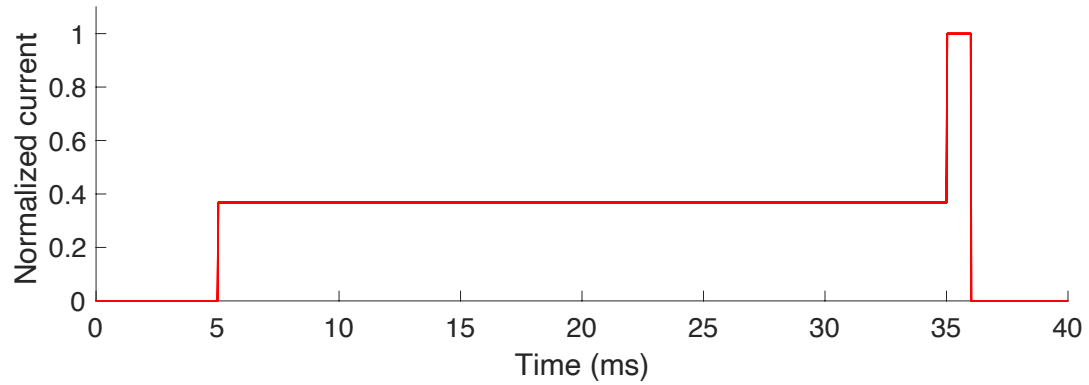
The figures to the left correspond to the activation thresholds estimates by the computational model for the $A\delta$ model when the Pin electrode is activating the nerve model. The activation thresholds were calculated when the fiber model was placed underneath the electrode in the middle of the epidermis ($A\delta$ model) or dermis ($A\beta$ model). For the $A\delta$ model the Na_P and K_A current were reduced to 10 % of its original value. The HCN current and the Na_{TTXs} current was increased by a factor of 2 and 8 respectively. The figures to the right correspond to the activation thresholds estimated by the computational model for the $A\beta$ model when the Patch electrode is activating the nerve model. For the $A\beta$ model the HCN current was reduced to 10 % of its original value and the Na_P density was increased by a factor of 1.5.



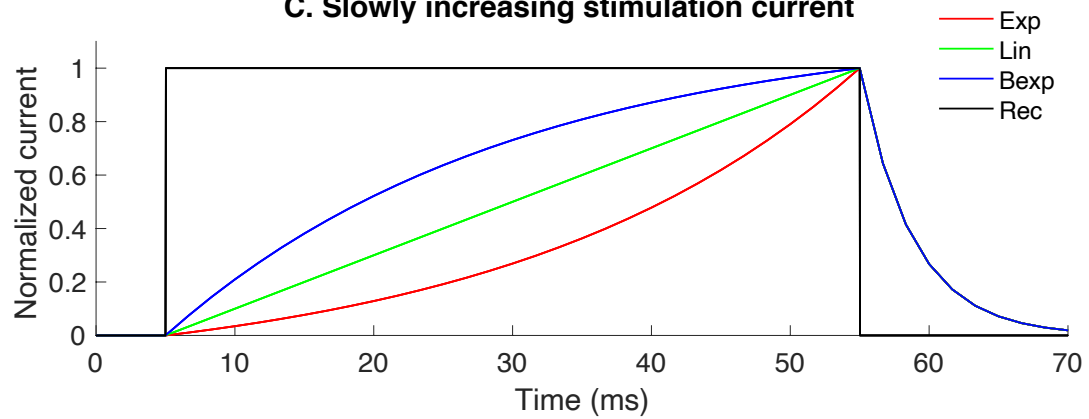
A. Strength duration curve

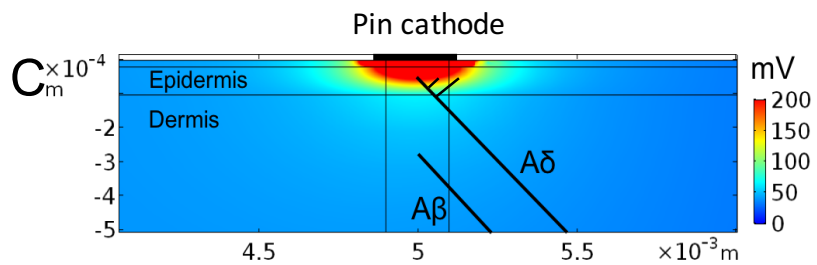
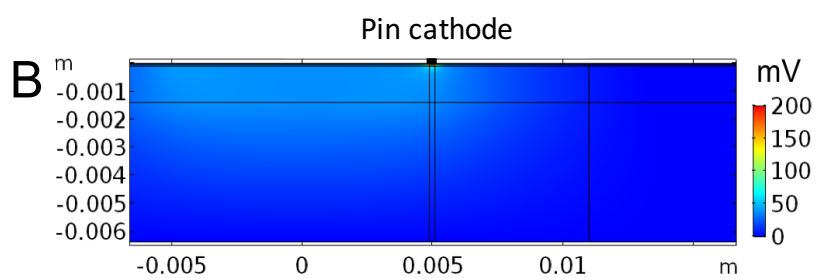
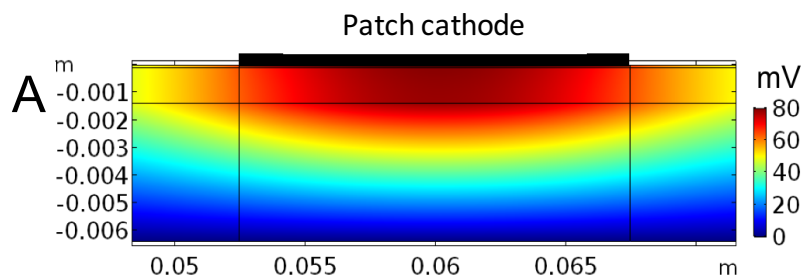


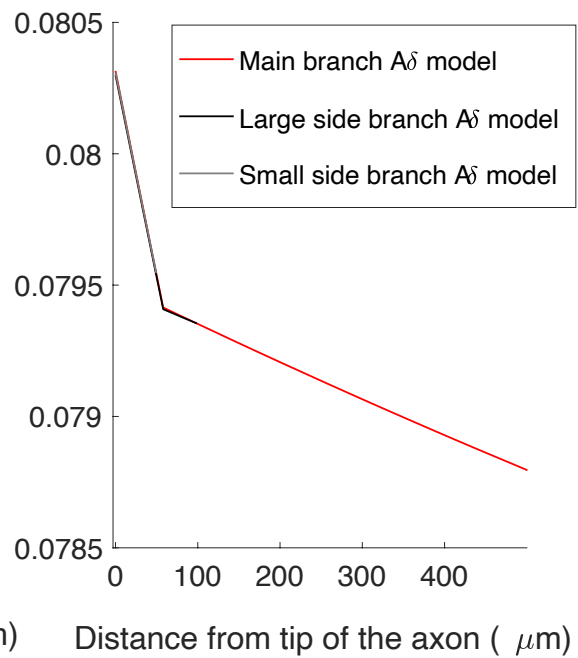
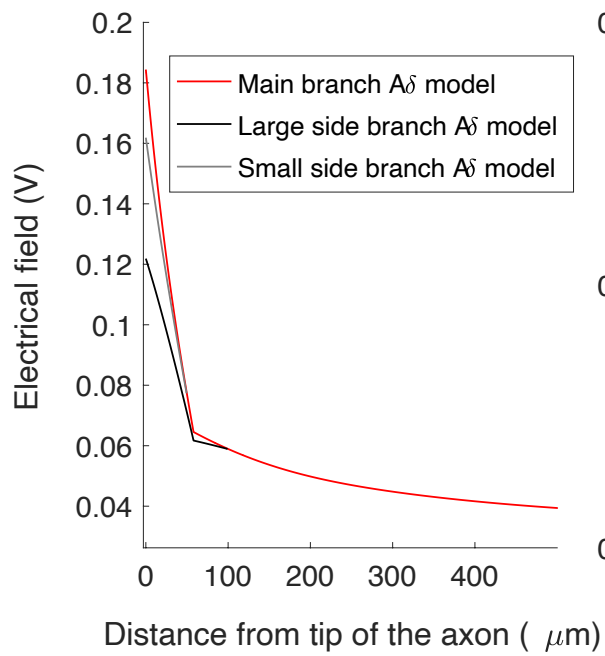
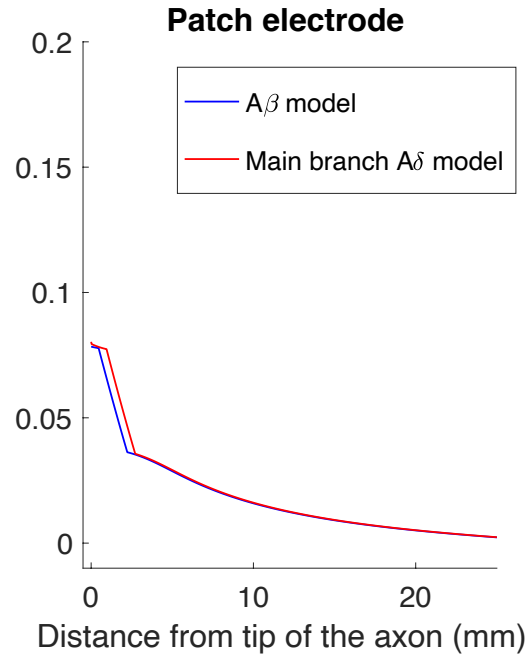
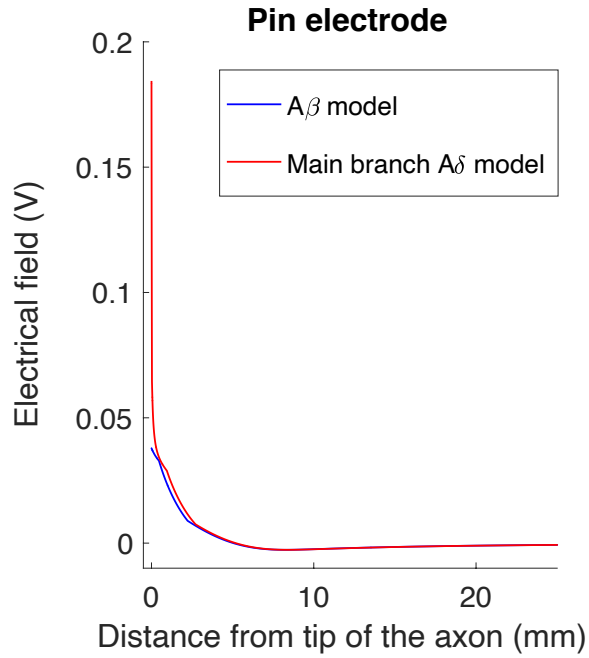
B. Threshold electrotonus

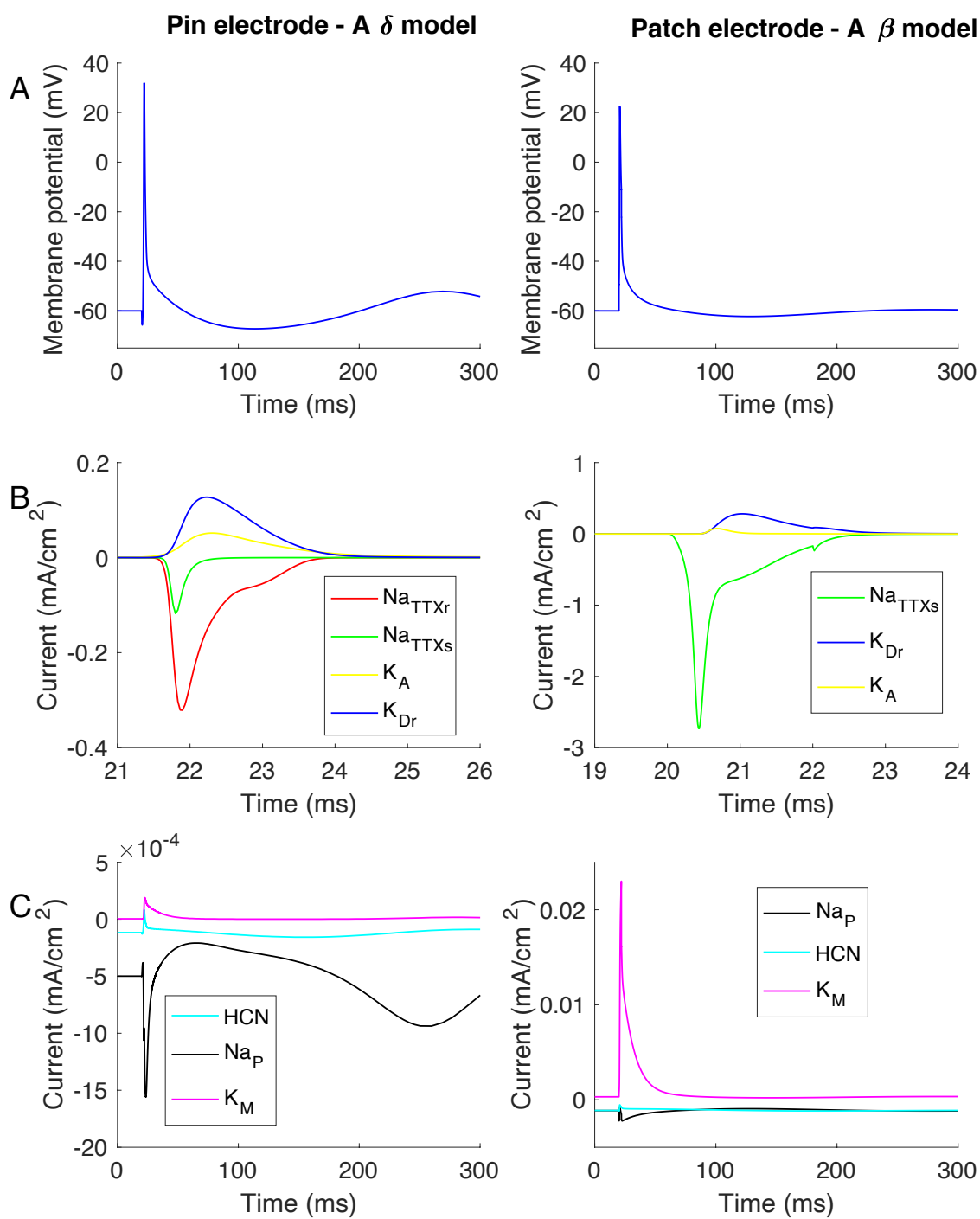


C. Slowly increasing stimulation current







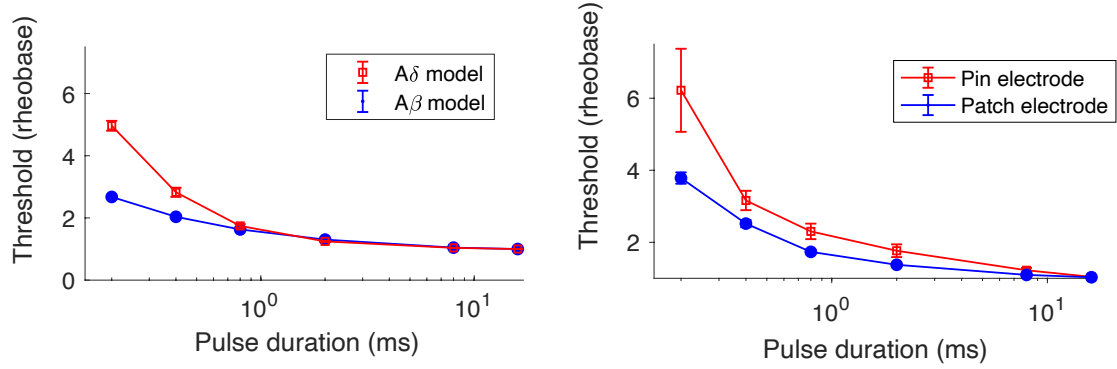


Computational model

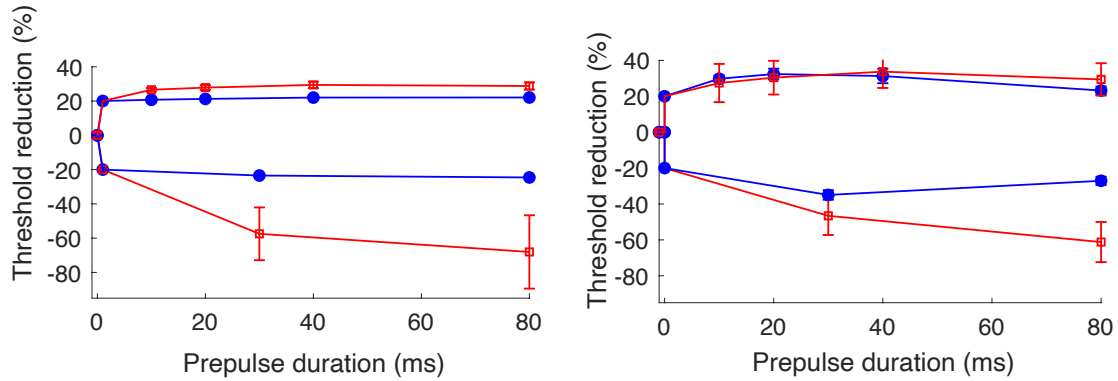
Experimental result

(18,20)

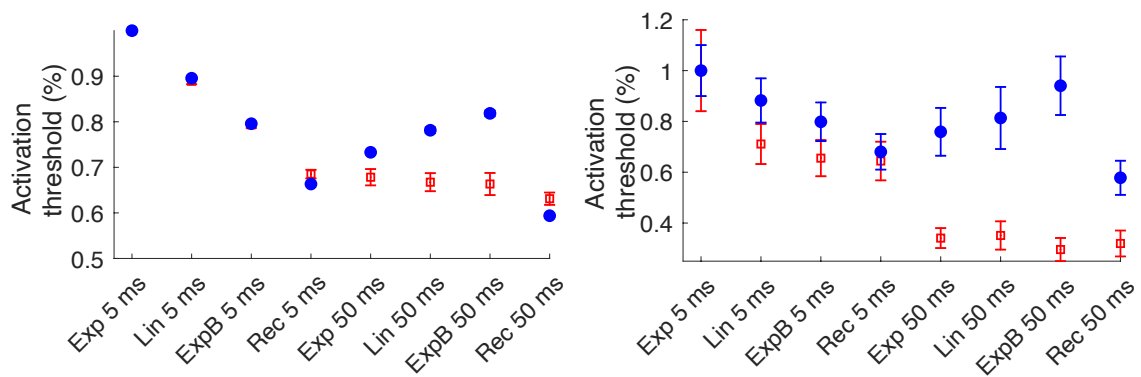
A. The strength duration curve

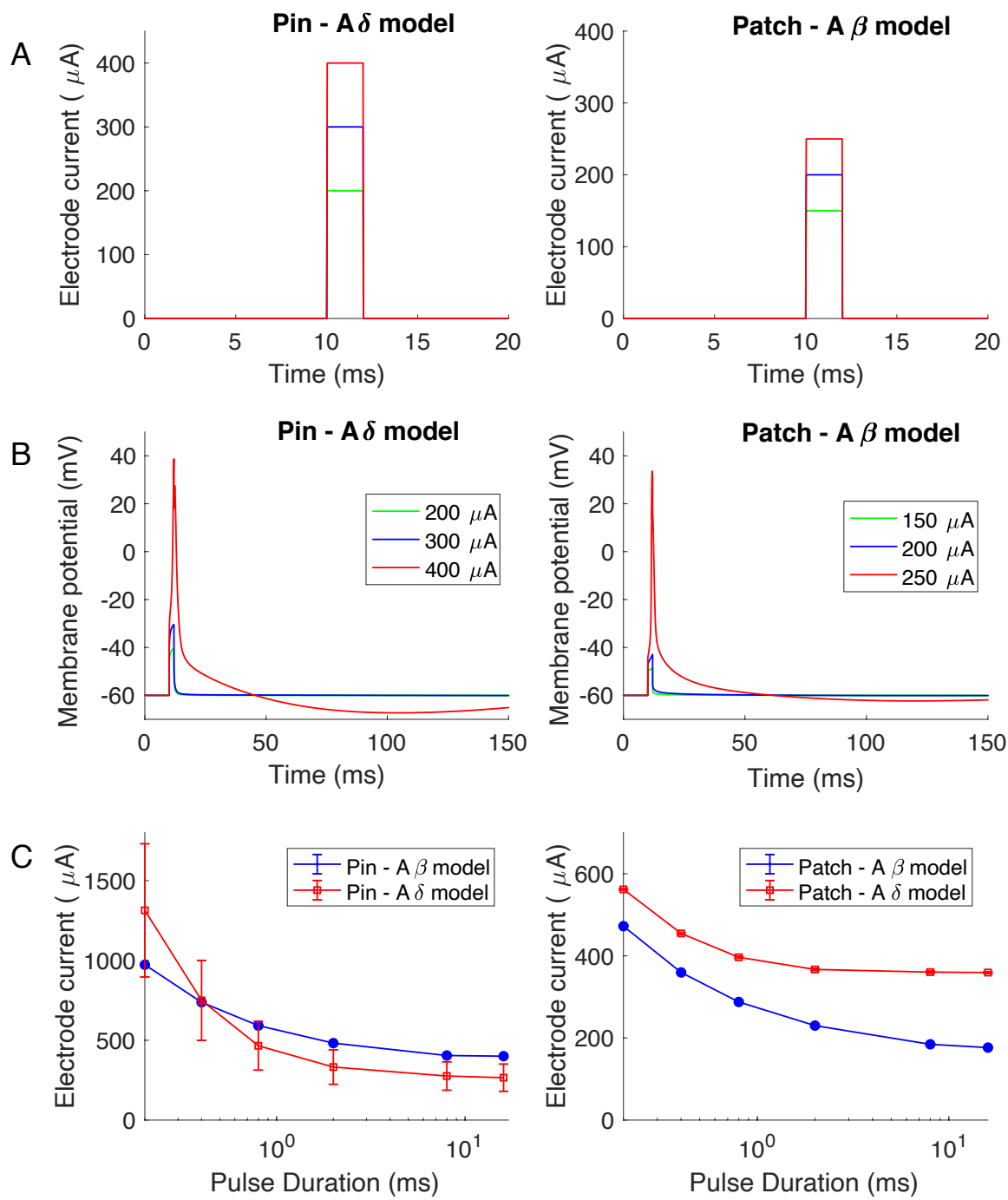


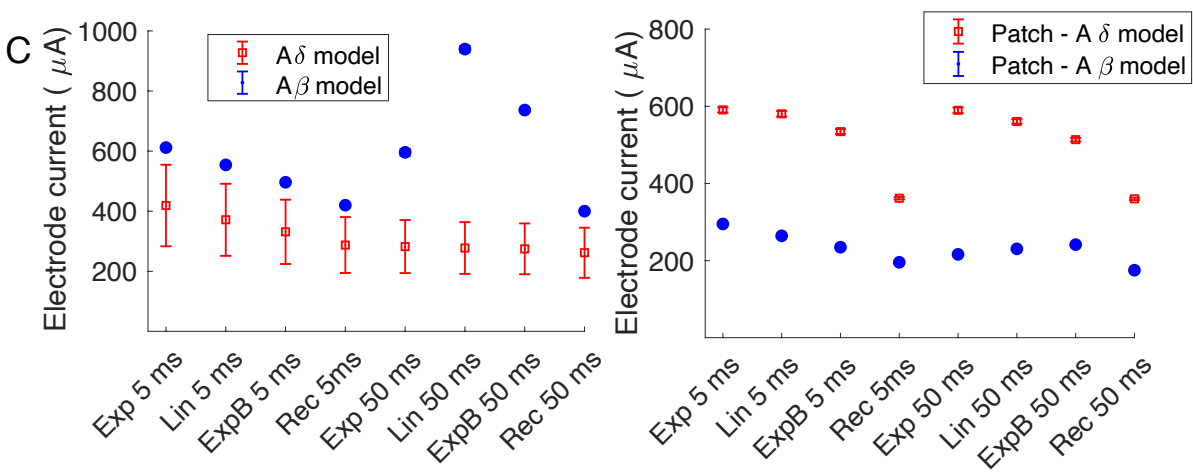
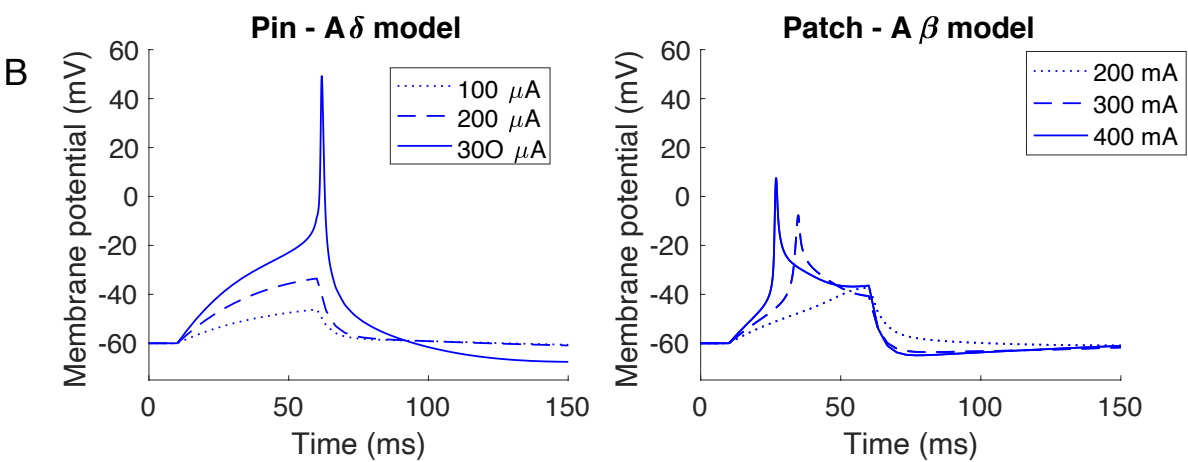
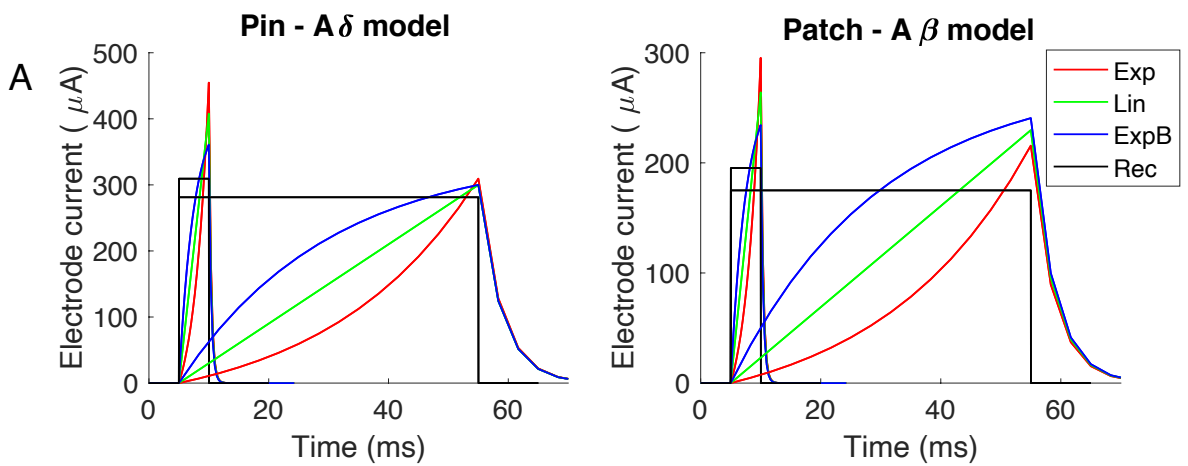
B. Threshold electrotonous



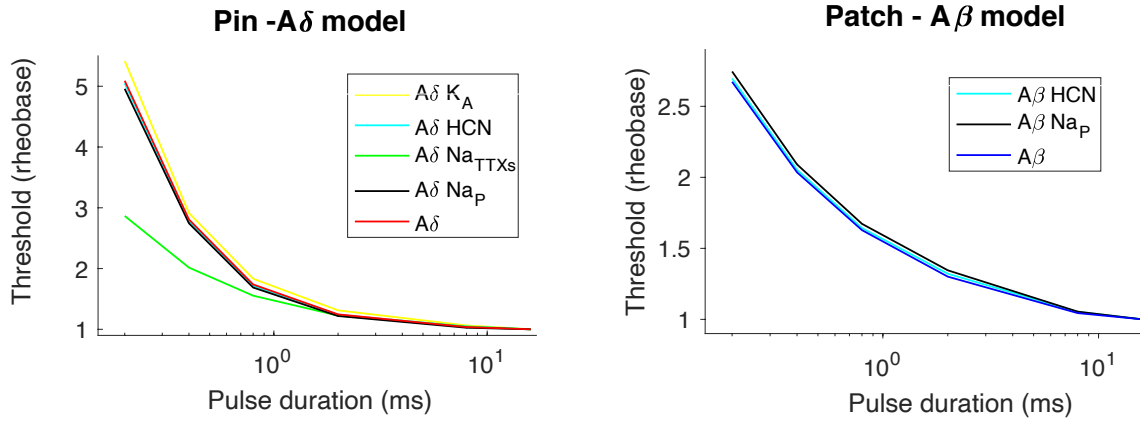
C. Slowly increasing stimulation current



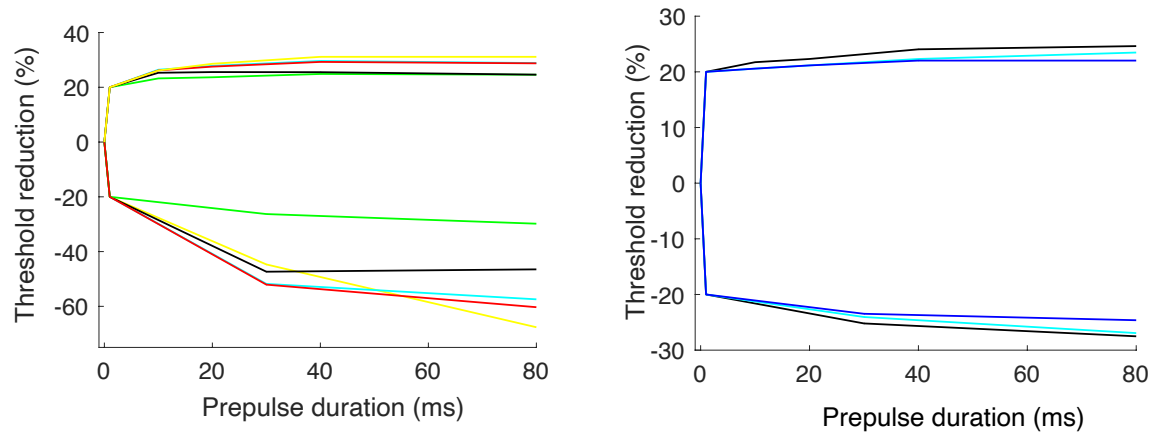




A. Strength duration curve



B. Threshold electrotonous



C. Slowly increasing stimulation current

